Prepared from FCVM PTO-1390 Transmittal Letter to the United States Designated/Elected Office (DQ/EO/US) JC10 Rec'd PCT/PTO 0.4 JAN 2002					
Customer No.	026418 10/030062				
Attorney's Docket No.:	GK-OEH-120 / 500814.20021				
U.S. Application No.:					
International Application No.:	PCT/DE00/02154				
International Filing Date:	JULY 4, 2000	4 JULY 2000			
Priority Date Claimed:	JULY 5, 1999	5 JULY 1999			
Title of Invention:	METHOD FOR THE MULTI-DIMENS PROTEOME	DD FOR THE MULTI-DIMENSIONAL ANALYSIS OF A			
Applicant(s) for (DO/EO/US):	Thomas MOORE and Anton HOR	N			
Applicant herewith submits to the United States Designated/Elected Office (DO/EO/US) the following items and other information: [X] 1. This is a FIRST submission of items concerning a filing under 35 U.S.C. 371. [] 2. This is a SECOND or SUBSEQUENT submission of items concerning a filing under 35 U.S.C. 371. [] 3. This express request to begin national examination procedures [35 U.S.C. 371 (f)] at any time rather than delay examination until the expiration of the applicable time limit set forth in 35 U.S.C. 371 (b) and PCT Articles 22 and [] 4. A proper Demand for International Preliminary Examination was made by the 19th month from the earliest claimed priority date. [X] 5. A copy of Publication No. WO 01/02848 11JAN01 the International Application as filed [35 U.S.C. 371(c)(2)] [A] a) is transmitted herewith (required only if not transmitted by the International Bureau) [B] b) has been transmitted by the international Bureau [C] c) is not required, as the application was filed in the United States Receiving Office (RO/US) [A] A translation of Publication No. WO 01/02848 11JAN01 the International Application into English [35 U.S.C. 371(c)(2)] [A] A mendments to the claims of the International Application under PCT Article 19 [35 U.S.C. 371(c)(3)] [A] D) have been transmitted by the International Bureau [B] D) have been made; however, the time limit for making such amendments has NOT expired. [B] D) have not been made and will not be made [B] A translation of the amendments to the claims under PCT Article 19 [35 U.S.C. 371(c)(3)] [B] An UNSIGNED Oath or declaration of the inventor(s) [35 U.S.C. 371(c)(4)] EXECUTED Decl/POA TO FOLLOW A translation of the annexes to the International Preliminary Examination Report under PCT Article 36 [35 U.S.C. 371(c)(5)]					
It 11. to 16. Below concern other document(s) or information included: It 11. An Information Disclosure Statement under 37 C.F.R. 1.97 and 1.98 An Assignment document for recording. A separate cover sheet (PTO-1619A) in compliance with 37 CFR 3.28 and 3.31 is included. It It It It It It It It					
EXPRESS MAIL No.: EL 915 668 453 US Deposited: January 4, 2002 I hereby certify that this correspondence is being deposited with the United States Postal Service Express mail under 37 CFR 1.10 on the date indicated above and is addressed to: BOX PCT, Commissioner for Patents, Washington, DC 20231. /Ruth Montalvo Date: January 4, 2002					

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D.C. 20231/

Ruth Montalvo

Date

Docket No.:GK-OEH-120/500814.20021

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applicant:

Thomas MOORE and Anton HORN

Serial No .:

Unknown (Int'l Appln. PCT/DE00/02154

filed July 4, 2000)

Filed:

Simultaneously herewith

For:

METHOD FOR THE MULTI-DIMENSIONAL ANALYSIS

OF A PROTEOME

PRELIMINARY AMENDMENT

Commissioner for Patents Washington, D.C. 20231

Sir:

Please amend the above-identified application, filed simultaneously herewith, as follows:

IN THE SPECIFICATION

Cancel the present specification and substitute therefor the enclosed substitute specification.

IN THE CLAIMS

Cancel claims 1-12 and add new claims 13-24, reading as follows:

--13. (New) A method for the multidimensional analysis of a proteome in which the biological material with the proteome to be analyzed is solubilized and the proteins belonging to the proteome are separated, quantitatively determined and identified, comprising the steps of:

subjecting the proteome to a number n of different separating processes for n>2 under standardized conditions in such a way that each of the liquid fractions m_1 obtained in a separating step supplies m_2 liquid fractions in a subsequent separating steps, wherein, after n separating steps, there are $m_1 * m_2 * \dots m_n = M$ liquid fractions;

identifying said $m_1 * m_2 * \dots m_n = M$ liquid fractions by τ different analysis processes qualitatively and/or quantitatively by identification processes, and determining said liquid ratio quantitatively by known quantification processes; and

after combining the analysis data, obtaining an n-dimensional image of the proteome which is characterized by identifiers and quantifiers and by the position in the n-dimensional data space.

- 14. (New) The method according to claim 13, wherein methods which separate according to the size of the protein and/or methods which separate according to the mass of the protein and/or methods which separate according to the charge of the protein and/or methods which separate according to the hydrophobicity of the protein and/or methods which separate according to the shape of the protein and/or methods which separate according to the shape of the protein and/or methods which separate according to the affinity of the protein, with respect to specific ligands, also to antibodies are selected as separating methods.
- 15. (New). The method according to claim 13, wherein methods for determining specific immunological characteristics and/or methods for determining specific catalytic activity and/or methods for determining chemical

modification of the proteins of the proteome are used as identification methods.

- 16. (New). The method according to claim 13, wherein methods for nonspecific determination of protein concentration with different sensitivities and/or quantitative determination methods for determining specific catalytic activities and/or quantitative immunological methods and/or quantitative binding assays are selected as quantification methods.
- 17. (New). The method according to claim 13, wherein the identification of individual proteins of the proteome is carried out directly by mass determination of the proteins.
- 18. (New) The method according to claim 13, wherein the identification of individual proteins is carried out according to protease digestion and mass identification of fragments.
- 19. (New) The method according to claim 13 wherein, after the separation step, the fractions are assembled in a two-dimensional multiple vessel system, in the manner of and with the layout of microtitration plates.
- 20. (New). The method according to claim 13 wherein, in the first separating step, the fractions are assembled in a defined grid, preferably in the n * 96 grid of microtitration technology.
- 21. (New) The method according to claim 13, wherein all identification and quantification steps are carried out in a defined grid, preferably in the n * 96 grid, with adaptable liquid handling technique.
- 22. (New) The method according to claim 21, wherein all identification steps and quantification steps are carried out with at least four two-dimensionally arranged, simultaneously working pipettor channels.
 - 23. (New). The method according to claim 13, wherein the first

dimension for separation is high-resolution size exclusion, ion exchange or hydrophobicity chromatography, which are known per se, in that the second dimension is carried out by parallel separation and fractionation of the fractions of the first dimension by a principle of separation other than that used for the first dimension, and in that each further separation and fractionation is carried out by parallel separating and fractionating methods with the fractions obtained from the preceding separating and fractionating steps.

24. (New) The method according to claim 13, wherein the analysis data for the n-dimensional image of the protein are assembled in a database.--

IN THE ABSTRACT OF THE DISCLOSURE

Cancel the present Abstract of the Disclosure and substitute therefor the enclosed Abstract of the Disclosure which is attached to the substitute specification..

REMARKS

Claims 1-12 have been cancelled and new claims 13-24 have been added. The amendments to the claims have been made only to improve the form of the claims for examination purposes.

The specification and abstract have been amended to conform it to U.S. format.

An early and favorable action on the merits is respectfully requested.

Respectfully submitted,

January 4, 2002 REED SMITH LLP 375 Park Avenue New York, NY 10152-1799 JEGforGHK:ram

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Substitute Specification

Abstract of the Disclosure

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IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Customer No.	026418		
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Applicant(s) for (DO/EO/US):	Thomas MOORE and Anton HORN		

SUBSTITUTE SPECIFICATION and ABSTRACT

Docket No.: GK-OEH-120/500814.20021

METHOD FOR THE MULTI-DIMENSIONAL ANALYSIS OF A PROTEOME

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CROSS-REFERENCE TO RELATED APPLICATIONS

This application claims priority of PCT Application Serial No. PCT/DE00/02154 filed July 4, 2000 and German Application No. 199 32 270.8 filed July 5, 1999, the complete disclosures of which are hereby incorporated by reference.

BACKGROUND OF THE INVENTON

Field of the Invention a)

The invention is directed to a method for the multidimensional analysis of a proteome in which the biological tissue with the proteome to be analyzed is solubilized and the proteins belonging to the proteome are separated, quantitatively determined and identified. The method is used in biochemistry, biotechnology, medicine and in the pharmaceutical industry for purposes including diagnostics and the development of biologically active substances. Special areas of use are opening up in fundamental research, e.g., for clarifying questions pertaining to developmental biology or cell differentiation and in related research for screening active ingredient banks, for the development and optimization of biologically active substances or for differentiating between normal and pathogenic states in organisms.

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Description of the Related Art b)

Recently, genomes of organisms have been sequenced completely or in large part [Fraser, C. M. et al.: The minimal gene complement of Mycoplasma genitalium, Science, 1995, Oct. 20, 270 (5235), 397-403; Fleischmann, R. D. et al.: Whole-genome random sequencing and assembly of Haemophilus influenzae Rd. Science, 1995, July 28, 269 (5223), 496-512; Blattner, F. R. et al.: The complete genome sequence of Escherichia coli K-12, Science, 1997, Sept. 5, 277 (5331), 1453-74; Goffeau, A. et al.: Life with 6000 genes, Science, 1996, Oct. 25, 274

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(5287), 546, 563-7]. Sequencing of cDNA portions has been even more intensive [Clark, M. S.: Comparative genomics: the key to understanding the Human Genome Project, Bioessays, 1999, Feb. 21 (2), 121-30; Evans, M. J. et al.: Gene trapping and functional genomics, Trends Genet., 1997, Sept. 13 (9), 370-4]. The sequence data are stored in databases. The clarification of the genome of an organism ultimately leads "only" to an understanding of the relatively static information content of the genetic material for this organism. With cDNA sequences, it is possible, in principle, to determine expression levels of the mRNA as they relate to specific cells and specific environments and accordingly to obtain a gene expression pattern of the RNA.

From a gene of the genome, it is possible a) to develop by different processes various mRNA types which code for divergent proteins, and b) to form a large number of extremely differently functioning proteins from this by means of posttranslational modification. Previously known modifications include phosphorylation and dephosphorylation, limited proteolysis, acetylation, methylation, adenylation, sulfation, glycosylation [McDonald, L. J., et al.: Enzymatic and nonenzymatic ADP-ribosylation of cysteins, Mol. Cell. Biochem., 1994 Sept., 138 (1-2), 221-6; Baenziger, J. U.: Protein-specific glycosyltransferases: how and why they do it!, FASEB J., 1994, Oct. 8 (13), 1019-25; Mimnaugh, E. G. et al.: The measurement of ubiquitin and ubiquitinated proteins, Electrophoresis, Feb. 1999, 20 (2), 418-28; Davis, P. J. et al.: Protein modification by thermal processing, Allergy, 1998, 53 (46 Suppl.), 102-5]. However, the expressed and modified proteins ultimately yield the pattern which describes the cell differentiation and the reaction to internal and external influences of cells. Most striking is the limited importance of knowing the genome for the realization of a defined biological state when the various cells in different organs and inside an organ are compared. For example, a liver paranchyma cell, a nerve cell of the brain and a mucosa cell of the intestine have the same set of genetic information but completely different functions brought about by the regulation of the expression of the genome in these cells and the regulation of the enzyme pattern and protein pattern within the cells and the

various tissues.

DNA

Static and descriptive, with exceptions

RNA

Transfer of information.

Quantity is regulated and transfers the information of the DNA to the protein plane.

Proteins

Maintaining cell structure, reaction to changes and signals.
Interactions with other cells. Quantity and

activity are regulated.

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The term "proteome" was first used in 1996 [Friedrich, G. A.: Moving beyond the genome projects, Nat. Biotechnol., Oct. 1996, 14 (10), 1234-7].

The proteome, that is, the totality of all proteins in a cell, with a definite development stage and under defined environmental conditions, is a much more dynamic representation of the physiological state of cells, organs and organisms. Proteome analysis investigates which parts of the genome are expressed and modified under defined, cell-specific conditions. This has led to rapidly growing interest in this field, leading to a growing number of publications (PubMed search term: Proteome; over the last 1 year: 64 hits; over the last 2 years: 99 hits; over the last 5 years: 122 hits), conferences and events on this subject.

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In order to obtain a quantifiable "picture" of a proteome, the following procedure is currently performed: In a first step, the biological materials must be solubilized and homogenized (exceptions: e.g., in a serum, they are in a homogenous solution). The proteins are isolated or separated in the second step and identified in the third step. In the fourth step, the obtained data are evaluated [Ben, R. H., et al.: Two dimensional electrophoresis, The state of the art and future directions, Proteome Research, New frontiers in functional genomics, Springer 1997, Chap. 2, 13-33].

1. Solubilization

Methods and arrangements known from biochemistry are used for this purpose, e.g., shear homogenizers, ultrasonic processing, high-pressure pressing. The difficulty consists in quantitative solubilization which does not destroy the function of the proteins as far as possible, because only quantitatively solubilized proteins provide a real picture of the specimen material in the subsequent second step (separation and detection of proteins) [Rabilloud, T.: Solubilization of proteins in 2-D electrophoresis, An outline, Methods Mol. Biol;., 1999, 112, 9-19; Rabilloud, T. et al.: Improvement of the solubilization of proteins in two-dimensional electrophoresis with immobilized pH gradients, Electrophoresis, Mar.-Apr. 1997, 18 (3-4), 307-16; Staudenmann, W. et al.: Sample Handling for proteome analysis, Electrophoresis, May 1998, 19 (6), 901-8].

2. Separation and detection

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At present, two-dimensional gel electrophoresis is essentially used for separating the proteins of the proteome. First tests with two-dimensional HPLC have been carried out. However, they have not yet achieved the separation effect of two-dimensional electrophoresis [Opiteck G. J. et al.: Comprehensive twodimensional high-performance liquid chromatography for the separation of overexpressed proteins and proteome mapping, Anal. Biochem. May 1998, 1; 258 (2): 349-61]. The first dimension of two-dimensional electrophoresis is isolation according to the isoelectric point, that is, ultimately, according to the charge characteristics of a protein. In the second dimension, the proteins are separated according to size in a denaturing sodium dodecyl sulfate gel. This separation technique has been known for about 20 years. An advantage of two-dimensional electrophoresis consists in the possibility of separating a relatively large number of proteins on a surface with high resolution. Presently, it is assumed that approximately 10,000 proteins can be detected in a two-dimensional gel of this kind [Klose, J. et al.: Two-dimensional electrophoresis of proteins: an updated protocol and implications for a functional analysis of the genome, Electrophoresis, 1995, June

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16 (6), 1034-59]. Another advantage is that it is possible to quantify the separated proteins by radioactive marking or after staining with techniques that are likewise known. These quantification methods are protein-specific, have a limited dynamic detection range, are generally difficult to automate and are dependent on the respective conditions of use (which often can not be reproduced) [James, P.: Of genomes and proteomes, Biochem. Biophys. Res. Commun., 1997, Feb. 3, 231 (1), 1-6]. They are only suitable for relative determinations. Quantification by immunological characteristics is problematic because blot techniques having limited meaningfulness in terms of quantitative information must be used for this purpose.

This results in a fingerprint-like pattern which characterizes the proteome.

This separation technique has the following disadvantages:

- limited dynamic range due to the load capacity of the separating gel
- the maximum quantity of proteins that may be used is limited to a range of μg to mg protein [James, P.: Of genomes and proteomes, Biochem. Biophys. Res. Commun., 1997, Feb. 3, 231 (1), 1-6]
- restriction of sample volume used
- separation is limited to two dimensions
- the ampholytes required for separation and the acrylamide gel material can lead to artifacts and can accordingly contribute to misinterpretations which are difficult to detect
- proteins that are present in very high concentrations result in relatively strong signals and overlap proteins in low concentrations, so that direct identification and quantification is impossible in this case
- the loss of the native conformation in denaturing separating gel causes the loss of biologically functional characteristics and impedes the identification of proteins by determining their biological characteristics, for example, their catalytic activity or specific bonding characteristics
- secondary analysis, such as the frequently used specific proteolysis of individual proteins, followed by determinations of mass necessitates a step for

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extracting from the gel or blot membrane which is difficult to automate.

3. Identification of proteins

Sequencing, mass analysis and estimation of the isoelectric point from the length of run in the gel and mass analysis of peptide fragments after separation from the gel and tryptic digestion in mass spectrometry are normally used for this purpose [Shevchenko, A. et al.: Linking genome and proteome by mass spectrometry: large-scale identification of yeast proteins from two-dimensional gels, Proc. Natl. Acad. Sci. USA, 1996, Dec. 10, 93 (25), 14440-5; Traini, M. et al.: Towards an automated approach for protein identification in proteome projects, Electrophoresis, 1998, Aug. 19 (11), 1941-9]. Features such as the catalytic activity of the proteins and the native conformation are almost completely excluded from the utilized separating technique and are not available for identification.

In particular, the known identification methods have the following advantages and disadvantages:

- The sequencing is carried out by Edman degradation in automated arrangements and is relatively costly and time-consuming. It requires greater quantities of the protein. Therefore, in spite of current further development for mass screening, it is less suitable [Gooley, A. A. et al.: A role for Edman degradation in proteome studies, Electrophoresis, 1997, June 18(7), 1068-72]. However, this analytic step is necessary in most cases for identification of primarily unknown proteins.
- The specificity of information of mass determination of a protein which should finally lead to its identification is increased in that the proteins undergo protease digestion after separation and the information obtained by means of mass analysis is compared with the masses of the peptide sequences predicted from the primary structure after tryptic digestion. Essentially two types of mass spectrometry are used: The first is Matrix Assisted Laser Desorption Ionization Mass Spectrometry (MALDI-MS) and the second is ElectroSpray Ionization Mass Spectrometry (ESI-MS) [Ducret, A. et al.: High Throughput protein characterization

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by automated reverse-phase chromatography/electrospray tandem mass spectrometry, Protein Sci., 1998, Mar 7 (3), 706-19; Parker, K. C. et al.: Identification of yeast proteins from two-dimensional gels: working out spot crosscontamination, Electrophoresis, 1998, Aug. 19 (11), 1920-32]. The first method has the advantage that it allows a very large mass range of up to 1 million Dalton to be analyzed and can be carried out in a relatively robust manner. However, it can be carried out only discontinuously. The ESI technique, on the other hand, can be appended almost continuously to separating techniques and is presently showing a sharp growth in the development of breadth of application as well as technological possibilities. The enormous advances achieved in recent years with both techniques allow mass resolutions to isotope distribution, that is, resolutions of less than 1 Dalton. In this way, a mass spectrum of peptide fragments is obtained according to sequence-specific, defined protease digestion or another defined splitting of the proteins. This spectrum is typical for every protein and is used for protein identification in sequence databases of proteins and expressed sequence tag banks. Since the identification of the protein by precise identification of the predicted peptides takes place after protease digestion, any posttranslational modification of the proteins, e.g., by glycosylation, interferes with detection. Further, fragmentation spectra of the individual peptides in the mass spectrometer can supply information about the amino acid sequence of the peptides. This sequence information can be used by itself or along with the other known protein data to identify this protein in a sequence database. This method of sequence analysis is not yet routinely used at present due to the difficulties of correct data interpretation. The limits of protein identification through mass spectrometry methods reside in the incomplete detection of all protein sequences in existing databases.

4. Data analysis

The characteristics of the individual detected proteins from separation in two-dimensional electrophoresis, such as quantity, isoelectric point and size, and the data for protein identification from additional steps, e.g., sequencing or mass

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spectrometry, are combined. This produces the picture of the totality of the proteins with their identity and quantity in the respective proteome.

OBJECT AND SUMMARY OF THE INVENTION

It is the primary object of the invention to improve and facilitate quantification and identification of the proteins of a proteome and to make it possible for certain proteins to be quantified and identified for the first time.

According to the invention, the proteins of the proteome are subjected to a number n of different separating processes under standardized conditions in such a way that each of the m_1 liquid fractions obtained in a separating step supplies m_2 liquid fractions in a subsequent separating step, wherein, after n separating steps, there are $m_1 * m_2 * \dots m_n = M$ liquid fractions which are identified by τ different analysis processes qualitatively and/or quantitatively by identification processes, known per se, and determined quantitatively by quantification processes which are likewise known per se, so that after combining the analysis data an n-dimensional image of the proteome is obtained which is characterized by identifiers and quantifiers and by the position in the n-dimensional data space.

The method according to the invention is not subject to the tight limitation on quantity due to the load capacity of previously used two-dimensional electrophoresis. Protein quantities in the range of several grams can be used. The separating matrices can be utilized repeatedly. In this way, greater reproducibility of results can be achieved. The sample material that is used is in liquid phase and is accordingly immediately accessible for subsequent analysis steps. The improved maintaining of native characteristics during separation makes possible analytic methods such as activity determination and immunological processes based on the native conformation of the analytes. The separation of analytes with the same charge characteristics and size characteristics is not possible in the two-dimensional electrophoresis that is usually used. However, this restriction is eliminated through the use of at least one further characteristic, such as the hydrophobicity of the analytes, for separation. After separation, the samples in fractions are also available

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for additional preparative tasks.

The invention will be described more fully in the following with reference to an embodiment example shown in the drawings.

BRIEF DESCRIPTION OF THE DRAWINGS

In the drawings:

Fig. 1 shows the separation of 1000 proteins in three dimensions. Figure 1 comprises:

Fig. 1a:

fractions 1 to 33

Fig. 1b:

fractions 33/34 to 67

Fig. 1c:

fractions 68 to 100;

Fig. 2 shows a graphic three-dimensional view of the fractions according to Fig. 1.

DESCRIPTION OF THE PREFERRED EMBODIMENTS

As an embodiment example, 1000 proteins are to be described by three characteristics A, B, C. These characteristics may be, e.g., size, charge and hydrophobicity. The characteristics are randomly distributed in the proteins. All proteins are numbered consecutively. Subsequently, separation is carried out according to characteristic A (e.g., size), resulting in 100 fractions a with the corresponding proteins. These fractions a are separated into 10 fractions b according to characteristic B (e.g., charge).

Each of these fractions b is subjected to separation based on characteristic C (e.g., hydrophobicity) and gives fractions c. In total, $100 \times 10 \times 10 = 10,000$ individual fractions are obtained. Each protein obtained by separation is uniquely allocated to one of the fractions a, b, c according to its characteristics. In the assignment according to Fig. 1, the respective fractions are designated by number. In this case, the fractions a are associated with characteristic A. They divide the possible value range of characteristic A into one hundred equal parts, i.e., assuming a value range from 0 to 100, value 1, for example, corresponds to range 0

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to 1, value 2 corresponds to range 1 to 2, ..., and value 100 corresponds to range 99 to 100. Analogously, the possible value ranges of characteristics B and C are divided into ten equal parts, i.e., value 1, for example, corresponds to range 1 to 10. On the average, every tenth fraction contains a protein.

Considered at random, there is a possibility of multiple assignments. In the example shown in the list according to Figs. 1a-c, there are 39 double occupancies and one triple occupancy of fractions.

For reasons of space and for the sake of clarity, the empty 9,000 fractions are not shown.

Fig. 1 contains the following list in tabular form:

Protein	Fractions	Fractions	Fractions
No.	a	b	С

Fig. 1a shows fractions a = 1 to 33, Fig. 2a shows fractions a = 33/34 to 67 and Fig. 1c shows fractions a = 68 to 100. Fig. 2 shows a three-dimensional diagram with the positions of the fractions occupied by proteins according to Fig. 1

While the foregoing description and drawings represent the present invention, it will be obvious to those skilled in the art that various changes may be made therein without departing from the true spirit and scope of the present invention.

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Assignment of Reference Numbers

A, B, C - characteristic of proteins

a, b, c - fraction

ABSTRACT OF THE DISCLOSURE

The invention is directed to a method for the multidimensional analysis of a proteome. The method is used in biochemistry, biotechnology, medicine and in the pharmaceutical industry for purposes including diagnostics and the development of biologically active substances. The object of the invention is to improve and facilitate quantification and identification of the proteins of a proteome and to make it possible for certain proteins to be quantified and identified for the first time. According to the invention, the proteins of the proteome are subjected to a number n of different separating processes under standardized conditions in such a way that each of the liquid fractions m_1 obtained in a separating step supplies m_2 liquid fractions in a subsequent separating step, wherein, after n separating steps, there are $m_1 * m_2 * \dots m_n = M$ liquid fractions which are identified by o different analysis processes qualitatively and/or quantitatively by identification processes, known per se, and determined quantitatively by quantification processes which are likewise known per se, so that, after combining the analysis data in a database, an n-dimensional image of the proteome is obtained which is characterized by identifiers and quantifiers and by the position in the n-dimensional data network.

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

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GK-OEH-119 / 500814.20021		
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JULY 4, 2000	4 JULY 2000	
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METHOD FOR THE MULTI-DIMENSIONAL ANALYSIS OF A PROTEOME		
Thomas MOORE and Anton HORN		
	GK-OEH-119 / 500814.20021 PCT/DE00/02154 JULY 4, 2000 JULY 5, 1999 METHOD FOR THE MULT PROTEOME	

MARKED-UP/BOLDED SUBSTITUTE SPECIFICATION and ABSTRACT

10/030062 531 Rec'd PC7 04 JAN 2002

-1-

Docket No.: GK-OEH-120/500814.20021

METHOD FOR THE MULTI-DIMENSIONAL ANALYSIS OF A PROTEOME

CROSS-REFERENCE TO RELATED APPLICATIONS

This application claims priority of PCT Application Serial

No. PCT/DE00/02154 filed July 4, 2000 and German Application No. 199 32

270.8 filed July 5, 1999, the complete disclosures of which are hereby incorporated by reference.

BACKGROUND OF THE INVENTON

a) Field of the Invention

The invention is directed to a method for the multidimensional analysis of a proteome in which the biological tissue with the proteome to be analyzed is solubilized and the proteins belonging to the proteome are separated, quantitatively determined and identified. The method is used in biochemistry, biotechnology, medicine and in the pharmaceutical industry for purposes including diagnostics and the development of biologically active substances. Special areas of use are opening up in fundamental research, e.g., for clarifying questions pertaining to developmental biology or cell differentiation and in related research for screening active ingredient banks, for the development and optimization of biologically active substances or for differentiating between normal and pathogenic states in organisms.

b) Description of the Related Art

Recently, genomes of organisms have been sequenced completely or in large part [Fraser, C. M. et al.: The minimal gene complement of Mycoplasma genitalium, Science, 1995, Oct. 20, 270 (5235), 397-403; Fleischmann, R. D. et al.: Whole-genome random sequencing and assembly of Haemophilus influenzae Rd. Science, 1995, July 28, 269 (5223), 496-512; Blattner, F. R. et al.: The complete genome sequence of Escherichia coli K-12, Science, 1997, Sept. 5, 277 (5331),

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1453-74; Goffeau, A. et al.: Life with 6000 genes, Science, 1996, Oct. 25, 274 (5287), 546, 563-7]. Sequencing of cDNA portions has been even more intensive [Clark, M. S.: Comparative genomics: the key to understanding the Human Genome Project, Bioessays, 1999, Feb. 21 (2), 121-30; Evans, M. J. et al.: Gene trapping and functional genomics, Trends Genet., 1997, Sept. 13 (9), 370-4]. The sequence data are stored in databases. The clarification of the genome of an organism ultimately leads "only" to an understanding of the relatively static information content of the genetic material for this organism. With cDNA sequences, it is possible, in principle, to determine expression levels of the mRNA as they relate to specific cells and specific environments and accordingly to obtain a gene expression pattern of the RNA.

From a gene of the genome, it is possible a) to develop by different processes various mRNA types which code for divergent proteins, and b) to form a large number of extremely differently functioning proteins from this by means of posttranslational modification. Previously known modifications include phosphorylation and dephosphorylation, limited proteolysis, acetylation, methylation, adenylation, sulfation, glycosylation [McDonald, L. J., et al.: Enzymatic and nonenzymatic ADP-ribosylation of cysteins, Mol. Cell. Biochem., 1994 Sept., 138 (1-2), 221-6; Baenziger, J. U.: Protein-specific glycosyltransferases: how and why they do it!, FASEB J., 1994, Oct. 8 (13), 1019-25; Mimnaugh, E. G. et al.: The measurement of ubiquitin and ubiquitinated proteins, Electrophoresis, Feb. 1999, 20 (2), 418-28; Davis, P. J. et al.: Protein modification by thermal processing, Allergy, 1998, 53 (46 Suppl.), 102-5]. However, the expressed and modified proteins ultimately yield the pattern which describes the cell differentiation and the reaction to internal and external influences of cells. Most striking is the limited importance of knowing the genome for the realization of a defined biological state when the various cells in different organs and inside an organ are compared. For example, a liver paranchyma cell, a nerve cell of the brain and a mucosa cell of the intestine have the same set of genetic information but completely different functions brought about by the regulation of the expression of the genome in these cells and the regulation of the enzyme pattern and protein pattern within the cells and the various tissues.

DNA	RNA	Proteins
Static and descriptive,	Transfer of information.	Maintaining cell structure,
with exceptions	Quantity is regulated and	reaction to changes and
	transfers the information of	signals.
	the DNA to the protein plane.	Interactions with other
		cells. Quantity and
		activity are regulated.

The term "proteome" was first used in 1996 [Friedrich, G. A.: Moving beyond the genome projects, Nat. Biotechnol., Oct. 1996, 14 (10), 1234-7].

The proteome, that is, the totality of all proteins in a cell, with a definite development stage and under defined environmental conditions, is a much more dynamic representation of the physiological state of cells, organs and organisms. Proteome analysis investigates which parts of the genome are expressed and modified under defined, cell-specific conditions. This has led to rapidly growing interest in this field, leading to a growing number of publications (PubMed search term: Proteome; over the last 1 year: 64 hits; over the last 2 years: 99 hits; over the last 5 years: 122 hits), conferences and events on this subject.

In order to obtain a quantifiable "picture" of a proteome, the following procedure is currently performed: In a first step, the biological materials must be solubilized and homogenized (exceptions: e.g., in a serum, they are in a homogenous solution). The proteins are isolated or separated in the second step and identified in the third step. In the fourth step, the obtained data are evaluated [Ben, R. H., et al.: Two dimensional electrophoresis, The state of the art and future

directions, Proteome Research, New frontiers in functional genomics, Springer 1997, Chap. 2, 13-33].

1. Solubilization

Methods and arrangements known from biochemistry are used for this purpose, e.g., shear homogenizers, ultrasonic processing, high-pressure pressing. The difficulty consists in quantitative solubilization which does not destroy the function of the proteins as far as possible, because only quantitatively solubilized proteins provide a real picture of the specimen material in the subsequent second step (separation and detection of proteins) [Rabilloud, T.: Solubilization of proteins in 2-D electrophoresis, An outline, Methods Mol. Biol;., 1999, 112, 9-19; Rabilloud, T. et al.: Improvement of the solubilization of proteins in two-dimensional electrophoresis with immobilized pH gradients, Electrophoresis, Mar.-Apr. 1997, 18 (3-4), 307-16; Staudenmann, W. et al.: Sample Handling for proteome analysis, Electrophoresis, May 1998, 19 (6), 901-8].

2. Separation and detection

At present, two-dimensional gel electrophoresis is essentially used for separating the proteins of the proteome. First tests with two-dimensional HPLC have been carried out. However, they have not yet achieved the separation effect of two-dimensional electrophoresis [Opiteck G. J. et al.: Comprehensive two-dimensional high-performance liquid chromatography for the separation of overexpressed proteins and proteome mapping, Anal. Biochem. May 1998, 1; 258 (2): 349-61]. The first dimension of two-dimensional electrophoresis is isolation according to the isoelectric point, that is, ultimately, according to the charge characteristics of a protein. In the second dimension, the proteins are separated according to size in a denaturing sodium dodecyl sulfate gel. This separation technique has been known for about 20 years. An advantage of two-dimensional electrophoresis consists in the possibility of separating a relatively large number of

proteins on a surface with high resolution. Presently, it is assumed that approximately 10,000 proteins can be detected in a two-dimensional gel of this kind [Klose, J. et al.: Two-dimensional electrophoresis of proteins: an updated protocol and implications for a functional analysis of the genome, Electrophoresis, 1995, June 16 (6), 1034-59]. Another advantage is that it is possible to quantify the separated proteins by radioactive marking or after staining with techniques that are likewise known. These quantification methods are protein-specific, have a limited dynamic detection range, are generally difficult to automate and are dependent on the respective conditions of use (which often can not be reproduced) [James, P.: Of genomes and proteomes, Biochem. Biophys. Res. Commun., 1997, Feb. 3, 231 (1), 1-6]. They are only suitable for relative determinations. Quantification by immunological characteristics is problematic because blot techniques having limited meaningfulness in terms of quantitative information must be used for this purpose.

This results in a fingerprint-like pattern which characterizes the proteome.

This separation technique has the following disadvantages:

- limited dynamic range due to the load capacity of the separating gel
- the maximum quantity of proteins that may be used is limited to a range of µg to mg protein [James, P.: Of genomes and proteomes, Biochem. Biophys. Res. Commun., 1997, Feb. 3, 231 (1), 1-6]
- restriction of sample volume used
- separation is limited to two dimensions
- the ampholytes required for separation and the acrylamide gel material can lead to artifacts and can accordingly contribute to misinterpretations which are difficult to detect
- proteins that are present in very high concentrations result in relatively strong signals and overlap proteins in low concentrations, so that direct identification and quantification is impossible in this case
- the loss of the native conformation in denaturing separating gel causes the

loss of biologically functional characteristics and impedes the identification of proteins by determining their biological characteristics, for example, their catalytic activity or specific bonding characteristics

- secondary analysis, such as the frequently used specific proteolysis of individual proteins, followed by determinations of mass necessitates a step for extracting from the gel or blot membrane which is difficult to automate.

3. Identification of proteins

Sequencing, mass analysis and estimation of the isoelectric point from the length of run in the gel and mass analysis of peptide fragments after separation from the gel and tryptic digestion in mass spectrometry are normally used for this purpose [Shevchenko, A. et al.: Linking genome and proteome by mass spectrometry: large-scale identification of yeast proteins from two-dimensional gels, Proc. Natl. Acad. Sci. USA, 1996, Dec. 10, 93 (25), 14440-5; Traini, M. et al.: Towards an automated approach for protein identification in proteome projects, Electrophoresis, 1998, Aug. 19 (11), 1941-9]. Features such as the catalytic activity of the proteins and the native conformation are almost completely excluded from the utilized separating technique and are not available for identification.

In particular, the known identification methods have the following advantages and disadvantages:

- The sequencing is carried out by Edman degradation in automated arrangements and is relatively costly and time-consuming. It requires greater quantities of the protein. Therefore, in spite of current further development for mass screening, it is less suitable [Gooley, A. A. et al.: A role for Edman degradation in proteome studies, Electrophoresis, 1997, June 18(7), 1068-72]. However, this analytic step is necessary in most cases for identification of primarily unknown proteins.
- The specificity of information of mass determination of a protein which should finally lead to its identification is increased in that the proteins undergo

protease digestion after separation and the information obtained by means of mass analysis is compared with the masses of the peptide sequences predicted from the primary structure after tryptic digestion. Essentially two types of mass spectrometry are used: The first is Matrix Assisted Laser Desorption Ionization Mass Spectrometry (MALDI-MS) and the second is ElectroSpray Ionization Mass Spectrometry (ESI-MS) [Ducret, A. et al.: High Throughput protein characterization by automated reverse-phase chromatography/electrospray tandem mass spectrometry, Protein Sci., 1998, Mar 7 (3), 706-19; Parker, K. C. et al.: Identification of yeast proteins from two-dimensional gels: working out spot crosscontamination, Electrophoresis, 1998, Aug. 19 (11), 1920-32]. The first method has the advantage that it allows a very large mass range of up to 1 million Dalton to be analyzed and can be carried out in a relatively robust manner. However, it can be carried out only discontinuously. The ESI technique, on the other hand, can be appended almost continuously to separating techniques and is presently showing a sharp growth in the development of breadth of application as well as technological possibilities. The enormous advances achieved in recent years with both techniques allow mass resolutions to isotope distribution, that is, resolutions of less than 1 Dalton. In this way, a mass spectrum of peptide fragments is obtained according to sequence-specific, defined protease digestion or another defined splitting of the proteins. This spectrum is typical for every protein and is used for protein identification in sequence databases of proteins and expressed sequence tag banks. Since the identification of the protein by precise identification of the predicted peptides takes place after protease digestion, any posttranslational modification of the proteins, e.g., by glycosylation, interferes with detection. Further, fragmentation spectra of the individual peptides in the mass spectrometer can supply information about the amino acid sequence of the peptides. This sequence information can be used by itself or along with the other known protein data to identify this protein in a sequence database. This method of sequence analysis is not yet routinely used at present due to the difficulties of correct data interpretation. The limits of protein

identification through mass spectrometry methods reside in the incomplete detection of all protein sequences in existing databases.

4. Data analysis

The characteristics of the individual detected proteins from separation in two-dimensional electrophoresis, such as quantity, isoelectric point and size, and the data for protein identification from additional steps, e.g., sequencing or mass spectrometry, are combined. This produces the picture of the totality of the proteins with their identity and quantity in the respective proteome.

OBJECT AND SUMMARY OF THE INVENTION

It is the <u>primary</u> object of the invention to improve and facilitate quantification and identification of the proteins of a proteome and to make it possible for certain proteins to be quantified and identified for the first time.

According to the invention, the proteins of the proteome are subjected to a number n of different separating processes under standardized conditions in such a way that each of the m_1 liquid fractions obtained in a separating step supplies m_2 liquid fractions in a subsequent separating step, wherein, after n separating steps, there are $m_1 * m_2 * \dots m_n = M$ liquid fractions which are identified by τ different analysis processes qualitatively and/or quantitatively by identification processes, known per se, and determined quantitatively by quantification processes which are likewise known per se, so that after combining the analysis data an n-dimensional image of the proteome is obtained which is characterized by identifiers and quantifiers and by the position in the n-dimensional data space.

[Advantageous embodiment forms of the method are set forth in the subclaims 2 to 12.]

The method according to the invention is not subject to the tight limitation on quantity due to the load capacity of previously used two-dimensional electrophoresis. Protein quantities in the range of several grams can be used. The

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separating matrices can be utilized repeatedly. In this way, greater reproducibility of results can be achieved. The sample material that is used is in liquid phase and is accordingly immediately accessible for subsequent analysis steps. The improved maintaining of native characteristics during separation makes possible analytic methods such as activity determination and immunological processes based on the native conformation of the analytes. The separation of analytes with the same charge characteristics and size characteristics is not possible in the two-dimensional electrophoresis that is usually used. However, this restriction is eliminated through the use of at least one further characteristic, such as the hydrophobicity of the analytes, for separation. After separation, the samples in fractions are also available for additional preparative tasks.

The invention will be described more fully in the following with reference to an embodiment example shown in the drawings.

BRIEF DESCRIPTION OF THE DRAWINGS

In the drawings:

Fig. 1 shows the separation of 1000 proteins in three dimensions.

Figure 1 comprises:

Fig. 1a: fractions 1 to 33

Fig. 1b [2a]: fractions 33/34 to 67

Fig. 1c [3a]: fractions 68 to 100;

Fig. 2 shows a graphic three-dimensional view of the fractions according to Fig. 1.

DESCRIPTION OF THE PREFERRED EMBODIMENTS

As an embodiment example, 1000 proteins are to be described by three characteristics A, B, C. These characteristics may be, e.g., size, charge and hydrophobicity. The characteristics are randomly distributed in the proteins. All proteins are numbered consecutively. Subsequently, separation is carried out

according to characteristic A (e.g., size), resulting in 100 fractions a with the corresponding proteins. These fractions a are separated into 10 fractions b according to characteristic B (e.g., charge).

Each of these fractions b is subjected to separation based on characteristic C (e.g., hydrophobicity) and gives fractions c. In total, $100 \times 10 \times 10 = 10,000$ individual fractions are obtained. Each protein obtained by separation is uniquely allocated to one of the fractions a, b, c according to its characteristics. In the assignment according to Fig. 1, the respective fractions are designated by number. In this case, the fractions a are associated with characteristic A. They divide the possible value range of characteristic A into one hundred equal parts, i.e., assuming a value range from 0 to 100, value 1, for example, corresponds to range 0 to 1, value 2 corresponds to range 1 to 2, ..., and value 100 corresponds to range 99 to 100. Analogously, the possible value ranges of characteristics B and C are divided into ten equal parts, i.e., value 1, for example, corresponds to range 1 to 10. On the average, every tenth fraction contains a protein.

Considered at random, there is a possibility of multiple assignments. In the example shown in the list according to Figs. 1a-c, there are 39 double occupancies and one triple occupancy of fractions.

For reasons of space and for the sake of clarity, the empty 9,000 fractions are not shown.

Fig. 1 contains the following list in tabular form:

Protein	Fractions	Fractions	Fractions
No.	a	ь	С

Fig. 1a shows fractions a = 1 to 33, Fig. 2a shows fractions a = 33/34 to 67 and Fig. 1c shows fractions a = 68 to 100. Fig. 2 shows a three-dimensional diagram with the positions of the fractions occupied by proteins according to Fig. 1

While the foregoing description and drawings represent the

present invention, it will be obvious to those skilled in the art that various changes may be made therein without departing from the true spirit and scope of the present invention.

Abstract of the Disclosure

The invention is directed to a method for the multidimensional analysis of a proteome. The method is used in biochemistry, biotechnology, medicine and in the pharmaceutical industry for purposes including diagnostics and the development of biologically active substances. It is the object of the invention to improve and facilitate quantification and identification of the proteins of a proteome and to make it possible for certain proteins to be quantified and identified for the first time. According to the invention, the proteins of the proteome are subjected to a number n of different separating processes under standardized conditions in such a way that each of the liquid fractions m₁ obtained in a separating step supplies m₂ liquid fractions in a subsequent separating step, wherein, after n separating steps, there are $m_1 * m_2 * \dots m_n = M$ liquid fractions which are identified by o different analysis processes qualitatively and/or quantitatively by identification processes, known per se, and determined quantitatively by quantification processes which are likewise known per se, so that, after combining the analysis data in a database, an ndimensional image of the proteome is obtained which is characterized by identifiers and quantifiers and by the position in the n-dimensional data network.

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Description of the Invention

METHOD FOR THE MULTI-DIMENSIONAL ANALYSIS OF A PROTEOME

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The invention is directed to a method for the multidimensional analysis of a proteome in which the biological tissue with the proteome to be analyzed is solubilized and the proteins belonging to the proteome are separated, quantitatively determined and identified. The method is used in biochemistry, biotechnology, medicine and in the pharmaceutical industry for purposes including diagnostics and the development of biologically active substances. Special areas of use are opening up in fundamental research, e.g., for clarifying questions pertaining to developmental biology or cell differentiation and in related research for screening active ingredient banks, for the development and optimization of biologically active substances or for differentiating between normal and pathogenic states in organisms.

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- the ampholytes required for separation and the acrylamide gel material can lead to artifacts and can accordingly contribute to misinterpretations which are difficult to detect
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Sequencing, mass analysis and estimation of the isoelectric point from the length of run in the gel and mass analysis of peptide fragments after separation from the gel and tryptic digestion in mass spectrometry are normally used for this purpose [Shevchenko, A. et al.: Linking genome and proteome by mass spectrometry: large-scale identification of yeast proteins from two-dimensional gels, Proc. Natl. Acad. Sci. USA, 1996, Dec. 10, 93 (25), 14440-5; Traini, M. et al.: Towards an automated approach for protein identification in proteome projects, Electrophoresis, 1998, Aug. 19 (11), 1941-9]. Features such as the catalytic activity of the proteins and the native conformation are almost completely excluded from the utilized separating technique and are not available for identification.

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The sequencing is carried out by Edman degradation in automated arrangements and is relatively costly and time-consuming. It requires greater quantities of the protein. Therefore, in spite of current further development for mass screening, it is less suitable [Gooley, A. A. et al.: A role for Edman degradation in proteome studies, Electrophoresis, 1997, June 18(7), 1068-72]. However, this

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present due to the difficulties of correct data interpretation. The limits of protein identification through mass spectrometry methods reside in the incomplete detection of all protein sequences in existing databases.

4. Data analysis

The characteristics of the individual detected proteins from separation in two-dimensional electrophoresis, such as quantity, isoelectric point and size, and the data for protein identification from additional steps, e.g., sequencing or mass spectrometry, are combined. This produces the picture of the totality of the proteins with their identity and quantity in the respective proteome.

It is the object of the invention to improve and facilitate quantification and identification of the proteins of a proteome and to make it possible for certain proteins to be quantified and identified for the first time.

According to the invention, the proteins of the proteome are subjected to a number n of different separating processes under standardized conditions in such a way that each of the m_1 liquid fractions obtained in a separating step supplies m_2 liquid fractions in a subsequent separating step, wherein, after n separating steps, there are $m_1 * m_2 * \dots m_n = M$ liquid fractions which are identified by τ different analysis processes qualitatively and/or quantitatively by identification processes, known per se, and determined quantitatively by quantification processes which are likewise known per se, so that after combining the analysis data an n-dimensional image of the proteome is obtained which is characterized by identifiers and quantifiers and by the position in the n-dimensional data space.

Advantageous embodiment forms of the method are set forth in the subclaims 2 to 12.

The method according to the invention is not subject to the tight limitation on quantity due to the load capacity of previously used two-dimensional electrophoresis. Protein quantities in the range of several grams can be used. The separating matrices can be utilized repeatedly. In this way, greater reproducibility of results can be achieved. The sample material that is used is in liquid phase and is accordingly immediately accessible for subsequent analysis steps. The improved maintaining of native characteristics during separation makes possible analytic

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The invention will be described more fully in the following with reference to an embodiment example shown in the drawing.

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Fig. 1 shows the separation of 1000 proteins in three dimensions

Fig. 1a:

fractions 1 to 33

Fig. 2a:

fractions 33/34 to 67

Fig. 3a:

fractions 68 to 100;

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Fig. 2 shows a graphic three-dimensional view of the fractions according to Fig. 1.

As an embodiment example, 1000 proteins are to be described by three characteristics A, B, C. These characteristics may be, e.g., size, charge and hydrophobicity. The characteristics are randomly distributed in the proteins. All proteins are numbered consecutively. Subsequently, separation is carried out according to characteristic A (e.g., size), resulting in 100 fractions a with the corresponding proteins. These fractions a are separated into 10 fractions b according to characteristic B (e.g., charge).

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Each of these fractions b is subjected to separation based on characteristic C (e.g., hydrophobicity) and gives fractions c. In total, $100 \times 10 \times 10 = 10,000$ individual fractions are obtained. Each protein obtained by separation is uniquely allocated to one of the fractions a, b, c according to its characteristics. In the assignment according to Fig. 1, the respective fractions are designated by number. In this case, the fractions a are associated with characteristic A. They divide the possible value range of characteristic A into one hundred equal parts, i.e., assuming a value range from 0 to 100, value 1, for example, corresponds to range 0

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to 1, value 2 corresponds to range 1 to 2, ..., and value 100 corresponds to range 99 to 100. Analogously, the possible value ranges of characteristics B and C are divided into ten equal parts, i.e., value 1, for example, corresponds to range 1 to 10. On the average, every tenth fraction contains a protein.

Considered at random, there is a possibility of multiple assignments. In the example shown in the list according to Figs. 1a-c, there are 39 double occupancies and one triple occupancy of fractions.

For reasons of space and for the sake of clarity, the empty 9,000 fractions are not shown.

Fig. 1 contains the following list in tabular form:

Protein	Fractions	Fractions	Fractions			
No.	a	b	c			

Fig. 1a shows fractions a = 1 to 33, Fig. 2a shows fractions a = 33/34 to 67 and Fig. 1c shows fractions a = 68 to 100. Fig. 2 shows a three-dimensional diagram with the positions of the fractions occupied by proteins according to Fig. 1.

Assignment of Reference Numbers

A, B, C - characteristic of proteins

a, b, c - fraction

Patent Claims

- which the biological material with the proteome to be analyzed is solubilized and the proteins belonging to the proteome are separated, quantitatively determined and identified, characterized in that the proteins of the proteome are subjected to a number n of different separating processes for n>2 under standardized conditions in such a way that each of the liquid fractions m_1 obtained in a separating step supplies m_2 liquid fractions in a subsequent separating step, wherein, after n separating steps, there are $m_1 * m_2 * \dots m_n = M$ liquid fractions which are identified by τ different analysis processes qualitatively and/or quantitatively by identification processes, known per se, and determined quantitatively by quantification processes which are likewise known per se, so that after combining the analysis data an n-dimensional image of the proteome is obtained which is characterized by identifiers and quantifiers and by the position in the n-dimensional data space.
- 2. Method according to claim 1, characterized in that methods which separate according to the size of the protein and/or methods which separate according to the mass of the protein and/or methods which separate according to the charge of the protein and/or methods which separate according to the hydrophobicity of the protein and/or methods which separate according to the shape of the protein and/or methods which separate according to the affinity of the protein, with respect to specific ligands, also to antibodies are selected as separating methods.
- 3. Method according to claim 1, characterized in that methods for determining specific immunological characteristics and/or methods for determining specific catalytic activity and/or methods for determining chemical modification of the proteins of the proteome are used as identification methods.
- 4. Method according to claim 1, characterized in that methods for nonspecific determination of protein concentration with different sensitivities

and/or quantitative determination methods for determining specific catalytic activities and/or quantitative immunological methods and/or quantitative binding assays are selected as quantification methods.

- 5. Method according to claim 1, characterized in that the identification of individual proteins of the proteome is carried out directly by mass determination of the proteins.
- 6. Method according to claim 1, characterized in that the identification of individual proteins is carried out according to protease digestion and mass identification of fragments.
- 7. Method according to claim 1, characterized in that after the separation step the fractions are assembled in a two-dimensional multiple vessel system, preferably in the manner of and with the layout of microtitration plates.
- 8. Method according to claim 1, characterized in that in the first separating step the fractions are assembled in a defined grid, preferably in the n * 96 grid of microtitration technology.
- 9. Method according to claim 1, characterized in that all identification and quantification steps are carried out in a defined grid, preferably in the n * 96 grid, with adaptable liquid handling technique.
- 10. Method according to claim 9, characterized in that all identification steps and quantification steps are carried out with at least four two-dimensionally arranged, simultaneously working pipettor channels.
- 11. Method according to claim 1, characterized in that the first dimension for separation is high-resolution size exclusion, ion exchange or hydrophobicity chromatography, which are known per se, in that the second dimension is carried out by parallel separation and fractionation of the fractions of

the first dimension by means of a principle of separation other than that used for the first dimension, and in that each further separation and fractionation is carried out by parallel separating and fractionating methods with the fractions obtained from the preceding separating and fractionating steps.

12. Method according to claim 1, characterized in that the analysis data for the n-dimensional image of the protein are assembled in a database.

Abstract

The invention is directed to a method for the multidimensional analysis of a proteome. The method is used in biochemistry, biotechnology, medicine and in the pharmaceutical industry for purposes including diagnostics and the development of biologically active substances. It is the object of the invention to improve and facilitate quantification and identification of the proteins of a proteome and to make it possible for certain proteins to be quantified and identified for the first time. According to the invention, the proteins of the proteome are subjected to a number n of different separating processes under standardized conditions in such a way that each of the liquid fractions m₁ obtained in a separating step supplies m, liquid fractions in a subsequent separating step, wherein, after n separating steps, there are $m_1 * m_2 * \dots m_n = M$ liquid fractions which are identified by o different analysis processes qualitatively and/or quantitatively by identification processes, known per se, and determined quantitatively by quantification processes which are likewise known per se, so that, after combining the analysis data in a database, an n-dimensional image of the proteome is obtained which is characterized by identifiers and quantifiers and by the position in the n-dimensional data network.

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348	1	6	2	239	8	1	6	525	15	1	10					82	259	29	2	4
459	1	6	4	649	8	1	9	263	15	2	4		795	21	4	4	19	29	4	3
156	1	8	6	106	8	2	1	181	15	2	10		23	21	6	5	×530	29	242	6
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618	2	2	10	972	9	3	5	596	15	7	7		695	22	3	1	685	29	8	7
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729	~	19		997			6	361		2	10		769	27	1	5	607	32	10	17
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832		10		142	_		5	436		13	10	1	415	27	3	6	83	33	3	1 1
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Fig. 1a

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332 34 3 4	44 41 5 3	485 48 6 54	473 55 8 1	919 62 4 10
636 34 4 6	812 41 7 8	915 48 7 5	266 55 8 4	342 62 7 3
		458 48 27 88	393 55 8 10	
121 34 4 8				460 62 7 8
998 34 5 1	343 41 10 8	138 481 275 88	320 55 9 1	378 62 7 10
355 34 5 8	911 42 1 6	120 48 7 10	276 55 9 6	94 62 10 7
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810 34 6 2	417 42 3 2	30 49 1 6	518 56 1 10	464 63 2 8
34 34 6 9	782 42 4 3	571 49 2 1	245 56 2 6	949 63 3 2
				
734 34 7 1	807 42 5 7	936 49 3 10	870 56 4 5	394 63 4 1
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164 34 9 4	168 42 8 1	775 49 8 4	673 56 5 6	683 63 6 4
157 34 10 8	857 42 8 9	421 49 8 6	182 56 5 10	
				
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47 35 4 2	302 43 2 3	577 50 6 7	873 56 9 6	76 63 8 9
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671 35 6 8	906 43 5 10	711 50 8 9	412 57 5 7	739 63 9 5
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Fig. 1b

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676	71	3	3	1 17	32	79	9	1		358	86	8	2	1	758	92	7	4		397	98	9
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13	71	18	12	4 ⊱	35	80	3	3	1	496	86	9	8	1	747	93	2	9		892	99	4
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334	71	8	5	4	384 772	80	5	10	1	659 943	87	1	8	1	301 433	93	5	7	-	215 28	99	7
489	71	10		1	47	80	6	3	1	894	87	2	5	1	861	93	6	6	-	930	99	8
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732	72	4	9		54	81	2	4]	243	87	7	8		2619	94	191	55		268	2100	*2
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392	72	9	3	→	206	81	4	10	1	836	87	8	2	1	134	94	1	9		910	100	4
92	72	13	15		372	81	7	12	4	942	87	9	6	1	357	94	1 1	10		309	100	4
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723	73	14	13	-1	791	81	8	4	1	937	88	1	3	1	617	94	2	10		784	100	8
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Fig. 1c

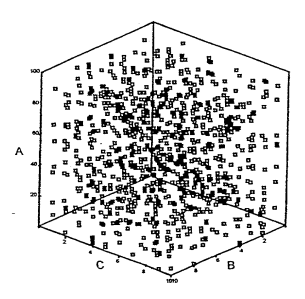


Fig. 2

UNITED STATES OF AMERICA COMBINED DECLARATION AND POWER OF ATTORNEY FOR PATENT APPLICATION

FILE NO. GK-OEH-120/ 500814.20021

below next to my name; that I verily believe that I am the original, first and sole inventor (if only one name is listed below) or a joint inventor (if plural inventors are named) of the subject matter which is claimed and for which a patent is sought on the invention entitled: METHOD FOR THE MULTI-DIMENSIONAL ANALYSIS OF A PROTEOME The specification of which is attached hereto. was filed on as United States patent application Serial Number was filed on July 4, 2000 as PCT international patent application No. PCT/DE00/02154 and was amended on (if any). X I hereby state that I have reviewed and understand the contents of the above identified specification, including the claims, as amended by any amendment referred to above. I acknowledge the duty to disclose all information known to be material to patentability in accordance with Title 37. Code of Federal Regulations, § 1.56. I hereby claim foreign priority benefits under Title 35, United States Code § 119 of any foreign application(s) for patent or inventor's certificate listed below and have also identified below any foreign application for patent or inventor's certificate having a filing date before that of the application on which priority is claimed: Prior Foreign Application(s) DATE OF FILING (day, month, year) PRIORITY COUNTRY APPLICATION NUMBER CLAIMED UNDER 35 U.S.C. § 119 199 32 270.8 YES x NO 05 July 1999 Germany I hereby appoint REED SMITH LLP and the members of the firm: Lloyd McAulay, Reg. No. 20,423; J. Harold Nissen, Reg. No. 17,283; Jules E. Goldberg, Reg. No. 24,408; Gerald H. Kiel, Reg. No. 25,116; Eugene LeDonne, Reg. No. 35,930; Stephen Chin, Reg. No. 39,938; Arthur Dresner, Reg. No. 24,403; Daniel Lent, Reg. No. 44,867; Samir R. Patel, Reg. No. 44,998; and Harry K. Ahn, Reg. No. 40,243, as attorneys with full power of substitution and revocation to prosecute all business in the Patent & Trademark Office connected therewith and to receive all correspondence. SEND CORRESPONDENCE TO: Gerald H. Kiel, Esa REED SMITH, LLP 375 Park Avenue New York, New York 10152-1799, U.S.A. DIRECT TELEPHONE CALLS TO: (212) 521-5400 I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or any patent issued thereon. INVENTOR'S SIGNATURE Ψ FULL NAME OF SOLE OR FIRST INVENTOR DATE Thomas MOORE 08.04.02 RESIDENCE COUNTRY OF CITIZENSHIP D-07751 Drackendorf, Germany Germany POST OFFICE ADDRESS Zur Laemmerlaide 15, D-07751 Drackendorf, Gelmany FULL NAME OF SECOND INVENTOR (IF ANY) INVENTOR'S SIGNATURE DATE Anton HORN RESIDENCE COUNTRY OF CITIZENSHIP D-07749 Jena, Germany Germany POST OFFICE ADDRESS Pennickental 4, D-07749 Jena, Germany

As a below named inventor, I hereby declare that: my residence, post office address and citizenship are as stated

X

COMBINED DECLARATION AND POWER OF FOR PATENT APPLICATION (continued)	ATTORNEY	File No. GK-OEH-119/ 500814.20021				
FULL NAME OF THIRD INVENTOR (IF ANY) Stefan KREUSCH	INVENTOR'S SIGNATURE		DATE 08.04.02			
RESIDENCE D-07743 Jena, Germany DE		COUNTRY O	F CITIZENSHIP			
POST OFFICE ADDRESS Freiligrathstrasse 90, D-07743 Jena, Ge	ermany					
FULL NAME OF FOURTH JOINT INVENTOR (IF ANY)		DATE				
RESIDENCE	COUNTRY O	F CITIZENSHIP				
POST OFFICE ADDRESS						
FULL NAME OF FIFTH INVENTOR (IF ANY)	INVENTOR'S SIGNATURE		DATE			
RESIDENCE		COUNTRY OF CITIZENSHIP				
POST OFFICE ADDRESS						
FULL NAME OF SIXTH JOINT INVENTOR (IF ANY)	INVENTOR'S SIGNATURE		DATE			
RESIDENCE		COUNTRY OF	F CITIZENSHIP			
PEST OFFICE ADDRESS						
FÜLL NAME OF SEVENTH JOINT INVENTOR (IF ANY)	INVENTOR'S SIGNATURE		DATE			
RESIDENCE		COUNTRY OF	- CITIZENSHIP			
POST OFFICE ADDRESS						
FULL NAME OF EIGHTH JOINT INVENTOR (IF ANY)	INVENTOR'S SIGNATURE		DATE			
RESIDENCE	COUNTRY OF CITIZENSHIP					
POST OFFICE ADDRESS						
FULL NAME OF NINTH JOINT INVENTOR (IF ANY)	INVENTOR'S SIGNATURE		DATE			
RESIDENCE		COUNTRY OF	CITIZENSHIP			
POST OFFICE ADDRESS						